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(54) Tissue scaffold anchor for cartilage repair

(57) A device (10) for attaching a tissue replacement careful of a box not (42) has a platform (12) positionable in substantially parallel relationship to the bone (42) for retaining the tissue scaffold proximate to the bone (42) for retaining the tissue scaffold proximate to the bone (42). A post (14) extends from the platform (12) and is insertable into a hole (40) formed in the bone (42). One or more ribs (16) extend from a side surface of the post (14) along a portion of its length. The sids (16) have an

increasing cross-sectional area to establish an increasing interference fit relative to the hole (40) in the bone tissue (42). The ribs (16) have a sharp edge that grips the sides of the hole (40) in the bone (42) such that the ribs (16) restrict rotation or withdrawal of the device (10).

Description

Field of the Invention

[0001] The present invention relates to scaffold fixation devices useful in articular cartilage repair and more specifically to a device for fastening an articular cartilage scaffold to underlying bone.

Background of the Invention

[0002] Articular cartilage is a tissue that covers the articulating surfaces between bones in joints, such as the knee or elbow, which is subject to catastrophic or repetitive stress injury. Various means have been proposed 15 to address such injuries including repair via tissue engineering. Tissue engineering is defined as the application of engineering disciplines to either maintain existing tissue structures or to enable new tissue growth. This engineering approach generally includes the delivery of 20 a tissue scaffold that serves as an architectural support onto which cells may attach, proliferate, and synthesize new tissue to repair a wound or defect. Surgical use of a tissue scaffold requires a fixation means to secure the scaffold to the bone beneath the wounded cartilage site. Secure fixation of the scaffold within the wound site is necessary for proper healing.

[0003] Frequently, scaffolds, prostheses and fasteners used in orthopedic applications are made from synthetic absorbable biocompatible polymers which are 30 well known in the art. Such polymers typically are used to manufacture medical devices which are implanted in body tissue and absorb over time. Synthetic, absorbable, biocompatible aliphatic polyesters include homopolymers, copolymers (random, block, segment- 35 ed and graft) of monomers such as glycolic acid, glycolide, lactic acid, lactide(d, l, meso and mixtures thereof), ε-caprolactone, trimethylene carbonate and p-dioxanone. Numerous U.S. Patents describe these polymers, including 5,431,679; 5,403,347; 5,314,989 40 5,431,679; 5,403,347; and 5,502,159. Devices made of an absorbable material have the advantage that they are absorbed by the body after healing has occurred. [0004] U.S. Patent 5,067,964 describes an articular cartilage repair piece which includes a backing layer of 45 non-woven, felted fibrous material which is either uncoated or covered by a coating of tough, pliable material. A number of means are disclosed for fastening the repair piece to the underlying bone. U.S. Patents 5,306,311 and 5,624,463 describe a prosthetic, resorb- 50 able articular cartilage and methods of its fabrication and insertion. U.S. Patent 5,713,374 describes an attachment method to hold a biomaterial in place until healing occurs, U.S. Patents 5,632,745 and 5,749,874 and 5,769,899 describe a bioabsorbable cartilage repair 55 system.

[0005] Articular joint loading is very complex, involving high compressive loads combined with high shear loads associated with sliding articulation of the opposing surfaces. A device implanted into the articular joint space must have sufficient strength to withstand these loads. Particularly important is that the device should be fixed in the underlying bone so that it cannot rotate or separate from the bone under the action of high shear loads in the joint space. U.S. Patent 5,748,974 baches that if vascular invasion and cellular migration is to be effected between the healthy tissue and the scaffold, means must be provided to preclude rotation of the scaffold relative to the fixation device, but does not describe a means of keeping the fixation device itself from rotating in relation to the surrounding issues or from pulling in relation to the surrounding issues or from pulling in relation to the surrounding issues or from pulling in relation to the surrounding issues or from pulling in relation to the surrounding issues or from pulling in relation to the surrounding issues or from pulling in relation to the surrounding issues or from pulling in the properties.

15 [0006] Accordingly, it would be advantageous to provide a scaffold fixation device which has a fixation means that engages the bone to prevent rotation and separation.

Summary of the Invention

out.

[0007] The limitations of prior art devices for attaching a tissue scaffold to bone tissue are overcome by the present invention which includes an attachment device having a pletform positionable in substantially parallel relationship to the bone tissue. A poet extends from the platform and is insertable into a hole formed in the bone tissue. A least one rib extends from a surface of the post along a portion of its length from a first point distal to the platform the platform. The rib has a cross-sectional area that increases along the length of the rib the direction from the first point and the first point of the first point of the second point and establishes an interference fit relative to the hole in the one tissue to prevent rotation or the device relative to

Brief Description of the Figures

100081

the hone tissue

Figure 1 is a side elevation view of a scaffold fixation device in accordance with an exemplary embodiment of the present invention;

Figure 2 is a perspective view of the device of Figure

Figure 3 is a side elevation view of the device of Figure 1 deployed in bone;

Figure 4 is an exploded view of a second exemplary embodiment of the present invention; Figure 5 is a side elevation view of the device of

Figure 4, assembled; and
Figure 6 is a side elevation view of the device of

Figure 6 is a side elevation view of the device of Figure 4 deployed in bone.

Detailed Description of the invention

100091 Figure 1 shows a scaffold fixation device 10 for fastening an articular cartilage scaffold to underlying bone. The device 10 has a scaffold attachment platform 12 with a post 14 extending therefrom at approximately 90°. Depending upon the application, this angular relationship may be varied. Vertical ribs 16 extend along a portion of the length of the post 14 and taper downwards in width and height as they extend from edge 17 to chamfered distal tip 18. The taper of vertical ribs 16 enhances the ability of the vertical ribs 16 to gradually cut into surrounding bone during insertion of scaffold fixation device 10 into an appropriately sized hole in a bone to which the device 10 is attached. While the ribs 16 15 shown are in the form of a longitudinally bisected, elongated cone, other tapering shapes could be employed. such as an elongated wedge with or without a knifeedge bevei.

[0010] Figure 2 shows a perspective view of scaffold 20 fixation device 10 showing perforations 20 in diskshaped platform 12 that allow fluid and cells to travel to and from the scaffold promoting cell proliferation and ingrowth. While six triangular perforations 20 are shown in FIG. 2, the perforations 20 can be any number, size 25 or shape, e.g., circular or trapezoidal and accordingly are not limited to the shape or arrangement shown in the figures. A guide wire channel 22 extends longitudinally through fixation device 10 along the axis of post 14. As is known in the art, a guide wire may be utilized 30 to assist in placing the device 10, viz, by inserting an end of a guide wire into a hole bored in a bone and then threading the device 10 over the guide wire, i.e., via channel 22, such that the post 14 enters the hole in the bone (See FIG. 3).

[0011] Figure 3 shows a side elevation view of scaffold fixation device 10 which has been surgically positioned within a hole 40 drilled in bone tissue 42. The diameter of the hole 40 is selected such that an interference fit is made between the hole 40 and post 14 with 40 vertical ribs 16. That is, hole 40 has diameter which is less than the outermost diameter of vertical ribs 16. Preferably, hole 40 has a diameter that is the same as or slightly smaller than the outermost diameter (root diameter) of post 14 (not including ribs 16). The scaffold fixation device 10 is preferably fabricated from a material that is sufficiently unyielding such that post 14 and vertical ribs 16 have sufficient radial stiffness and strength to cause the vertical ribs 16 to cut into the bone tissue 42 surrounding the hole 40. This intrusion into the bone 50 42 has the effect of rotationally fixing the scaffold fixation device 10 to the bone tissue 42. In addition, axial fixation of the device 10 is achieved by vertical ribs 16, the sharp edges 17 of which engage trabecular bone tissue 42 when subjected to an axial force which would otherwise 55 pull the scaffold fixation device 10 out of the hole 40 in the bone 42. A hole 44 is drilled in cartilage tissue 46 with a diameter at least as large as the outermost diameter of platform 12 to accommodate the platform 12 therein in a position permitting the scaffold 47 (shown diagrammatically in dotted lines and displaced slightly) to be attached to the device 10 by sutures or adhesives,

5 in a known manner. The depths of hole 40 in the bone and the hole 44 in the cartilage are selected such that, when post 14 is inserted completely into hole 40, upper surface 50 of platform 12 is in alignment with or slightly below upper surface 52 of the bone tissue 42, i.e., the platform 12 may be countersuris into the bone 42. The

19 platform 12 may be countersumk into the bone 42. The scaffield 47 is accommodated within hole 44 in the cartillage (between platform 12 and upper cartillage surface 54). Post 14 may also have a chamfered lower edge 16 which aids in guiding post 14 into the hole 40 in the bone (18sue 42. As noted above, a surgical guide wire may be passed through guide wire channel 22 during surgery to align scaffield fixation device 10 with bone hole 40. The fixation device 10 may be made from a non-procus amterial or from materials that are partially or wholly porcus.

to allow cell invasion into the device.

[0012] A two-piece embodiment of the invention is shown in Figures 4 through 6, which show a two-piece scaffold fixation device 130 similar to a device described in EP-A-1 129 675.

25 [0013] Figures 14 through 20 and the associated description thereof being particularly relevant in describing the interlocking relationship displayed by a two-piece scaffold fixation device.

[0014] Figure 4 shows a two-piece scaffold fixation of device 130 with top component 132 and fixation component 134. The top component 132 has a scaffold attachment platform 112 from which extends a coupling pin 114 with a pair of latches 115, 118 projecting from corresponding resilient arms 120, 122. The coupling pin 35 114 telescopes into a mating axial bore 124 in the fixa-

114 (usescopes into a missing assal ore 124 in the insation component 134, with the latches 115, 115 clipping over an Internal ledge 126 when the pin 114 is pressed fully home into the bore 124. The fixation component 124 has vertical ancioning rise 150 having a similar form 124 has vertical ancioning rise 150 having a similar form 124 has vertical ancioning rise 150 having a similar form 124 have 125 have 12

[0015] Figure 6 shows catchied fixation device 130 after having been surgically inserted in bone issue 162, showing the vertical anchoring rish 180 embedded in the bone issue 162 surrounding hole 160 to prevent rotation of fixation component 134 within the hole 160. The device 130 would be utilized for attaching a scaffold (see FiG. 3) to a bone 162 by boring a suitable hole 161 in the bone 162. The fixation component 134 is inserted into the hole 160 and driven home. The coupling pin 114 of the top component 132 can then be inserted into boxe 124 of the fixation component and pressed in until the latches 116, 118 latch over ledge 126 (See FiG. 4).

[0016] Although Fizures 1-5 show a certain number

[0017] Suitable materials from which the scafflot fixation device 10, 130 may be formed include biocompatation device 10, 130 may be formed include biocompatble polymers such as alliphatic polyesters, polyporthossters, polyamides and polyaliyene oxides. The present inpolyamides and polyaliyene oxides. The present inpation also can be formed from absorbable polymers, glasses or ceramics compation actious phosphatic and other biocompatible metal caddes (i.e., CaO), metals, combinations of metals, autograft, allograft, or xenograft bone listone.

[0018] In the preferred embodiment, the scaffold fix- 20 ation device 10, 130 is formed from aliphatic polymer and copolymer polyesters and blends thereof. The aliphatic polyesters are typically synthesized in a ring opening polymerization. Suitable monomers include but are not limited to lectic acid, lactide (including L-, D-, meso and D,L mixtures), glycolic acid, glycolide, εcaprolactone, p-dioxanone (1,4-dioxan-2-one), trimethylene carbonate (1,3-dioxen-2-one), delta-valerolactone, beta-butyrolactone. epsilon-decalactone. 2,5-diketomorpholine, pivalolactone, alpha, alpha-diethylpropiolactone, ethylene carbonate, ethylene oxalate, 3-methyl-1,4-dioxane-2,5-dione, 3,3-dlethyl-1,4-dioxan-2,5-dione, gamma-butyrolectone, 1.4-dioxepan-2-one, 1,5-dioxepan-2-one, 6,6-dimethyl-dioxepen-2-one, 6,8-dioxabicycloctane-7-one and combinations thereof. These monomers generally are polymerized in the presence of an organometallic catalyst and an initiator at elevated temperetures. The organometallic catalyst is preferably tin based, e.g., stannous octoate, end is present in the monomer mixture at a molar 40 ratio of monomer to catalyst ranging from about 10,000/1 to about 100,000/1. The initiator is typically an alkanol (including diols and polyols), a glycol, a hydroxyacid, or an amine, and is present in the monomer mixture at a molar ratio of monomer to initiator ranging from 45 about 100/1 to about 5000/1. The polymerization typically is carried out at a temperature range from about 80°C to about 240°C, preferably from about 100°C to about 220°C, until the desired molecular weight and viscosity are achieved.

[0019] In another embodiment of the present invention, the polymers and blends from which it is formed can be used as a therapeutic agent release martix. Prior to forming the device 10, 130, the polymer would be mixed with a therapeutic agent. The variety of different of therapeutic agents that can be used in conjunction with the polymers of the present invention is vast. In general, therapeutic agents which may be administered via the

pharmaceutical compositions of the invention include, without limitation: antiinfectives such as antibiotics and antiviral agents; chemotherapeutic agents (i.e. anticancer agents); anti-rejection agents; analgesics and analgesic combinations; anti-inflammatory agents; hor-

gesic combinations; anti-inflammatory agents; hormones such as steroids; growth factors, including bone morphogenic proteins (i.e. MPP's 1-7), bone morphogenic-like proteins (i.e. GFD-5, GFD-7 and GFD-8), epidermal growth factor (GEGF, fibroblast growth factor (i.e. FGF 1-9), platelet derived growth factor (PDGF), in-

sulin ike growth factor (IGF-I and IGF-In), transforming growth factor (KEGF); and other naturally derived or periodically engineered proteins, objectscharfeles, glyco-proteins, or lipoproteins. The foregoing growth factor are known to hose with skill in the art and described in The Cellular and Molecular Basis of Bone Formation and Registry Vicil Rosen and K. Scott Thies, published by R.G. Landes Company hereby incorporated herein by reference.

of interestication. Of which was a simple of the present invertion may be formulated by mixing one or more therapsutic agents with the polymer. Alternative, a therapsutic agent could be coated on to the polymer, preferebly with e pharmaceutical carrier can be used that does not dissolve the polymer. The therapsutic agent may be present as a liquid, a finely divided solid, or any other appropriate physical form. Typically, but optionely, the metrix will include one or more additives, such as diluents, carriers, excipients, stabilizers or the like.

[9021] The amount of therapeutic agent will depend on the particular drug being employed and medical condition being treated. Typicelly, the amount of drug represents about 0.001 percent to about 7.0 percent, more typically about 0.001 percent to about 7.0 percent, more typically about 0.001 percent to about 5.0 percent, more typically about 0.001 percent to about 5.0 percent, more typically about 0.001 percent to about 5.0 percent by weight of the matrix. The quantity and type of polymer incorporated into the drug delivery metrix will very depending on the release profile desired and the amount of drug employed.

[00:22] Upon contact with body fulids, the polymer undergoes gradual degradation (mainly through high youngst) with concomitant release of the dispersed drug for a sustained or strained period. This can result in prolonged delivery (over, say I to 5,000 hours, preferably 2 to 800 hours) of effective amounts (say, 0,000 millions) to 10 800 hours) of effective amounts (say) of the order nan be administered as is necessary depending on the 50 subject being treated, the severity of the affiction, the judgment of the prescribing physician, and the like. Following this or similar procedures, those skilled in the art will be able to prepare a variety of formulations.

Claims

1. A device for attaching a tissue scaffold to bone tis-

sue, comprising:

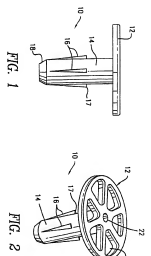
- a platform positionable in substantially parallel relationship to the bone tissue for retaining the tissue scaffold proximate to the bone tissue; a post extending from the platform, said post insertable into a hole formed in the bone tissue; at least one rib extending from a surface of said post along a portion of the length of said post from a first point distal to said platform to a second point intermediate said first point and said platform, said at least one rib having a crosssectional area that increases along the length of sald rib in the direction from said first point to said second point, said at least one rib es- 15 tablishing an interference fit relative to the hole in the bone tissue to prevent rotation of said device relative to the bone tissue.
- is shorter in length than the depth of the hole in the bone tissue and has a diverging surface that diverges from the surface of said post in the direction from said first point to said second point, said at least one rib terminating proximate said second point in a 25 sharp edge, said edge positionable within said hole for gripping the bone tissue to resist withdrawal of said device from the hole.
- 3. The device of Cleim 2, wherein said rib expands in 30 width in the direction from said first point to said second point.
- 4. The device of Claim 3, wherein said at least one rib has e shepe approximating a portion of an elongat- 35 ed cone.
- 5. The device of Claim 4, wherein said at least one rib is a plurality of ribs disposed about the outer peripheral surface of seid post.
- 6. The device of Claim 5, wherein said post has a chamfered end to aid in the introduction of said post into the hole in the hone tissue
- 7. The device of Claim 6, wherein said device has a guide wire hole extending axially there through to permit said device to be slipped over a guide wire having one end thereof positioned in the hole for guiding said device into the hole.
- 8. The device of Claim 1, wherein said post extends from said platform at about 90 degrees.
- 9. The device of Claim 2, wherein said platform has a 55 perforation therein to allow fluid and cell transmission through said perforation.

- 10. The device of Claim 2, wherein said device is formed from a material selected from the group consisting of biocompatible polymers, absorbable polymers, glasses, ceramics, metal oxides, bone tissue and therapeutic agents, alone or in combination
- 11. The device of Claim 1, wherein said platform and said post are monolithically formed.
- 12. The device of Claim 1, wherein said platform and said post are independently formed elements.
 - 13. The device of Claim 12, wherein said platform has a coupling pin extending from a surface thereof, said post having a hollow therein for matingly receiving said coupling pin to couple said platform and said post.
- 2. The device of Claim 1, wherein said at least one rib 20 14. The device of Claim 13, further including a latch resiliently mounted on said coupling pin, said latch having a locking position and a withdrawn position, said latch permitting said coupling pin to be inserted into said hollow in said post in a coupling relationship with said post when in said withdrawn position and retaining said coupling pin coupled to seid post in said locking position.
 - 15. A device for attaching a tissue scaffold to bone tissue, comprisino:
 - attaching means for attaching the scaffold to said device: retaining means coupled to said attaching
 - means for retaining said attaching means in proximity to the bone tissue, seid retaining means inserting into a hole formed in the bone tissue, said retaining means including gripping means extending from the outer periphery thereof for gripping the bone tissue proximate the hole to restrain seid device from rotating or pulling away from the bone tissue.
 - 16. The device of Claim 15, wherein said gripping means have a gradually increasing interference fit relative to the hole in the hone tissue
 - 17. The device of Claim 16, wherein said gripping means are fully insertable within the hole in the bone tissue and have a sharp edge which gouges into the bone tissue proximate the hole.
 - 18. The device of Claim 17, wherein said attaching means has apertures therein for allowing the flow of fluids and cells there through.
 - 19. The device of Claim 18, further including coupling means for coupling said attaching means and said

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retaining means.

20. The device of Claim 17 wherein said attaching means and said retaining means are formed from a composition with a porosity permitting ingrowth of 5 cells.



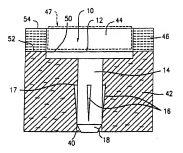
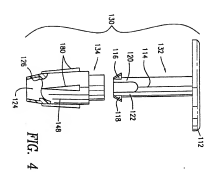
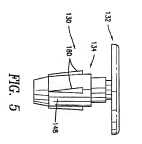


FIG. 3





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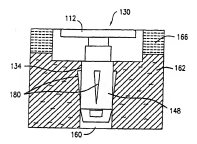


FIG. 6